SINCE FILE

ENTRY

0.21

TOTAL

SESSION 0.21 10/764,389

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\*\*\* YOU HAVE NEW MAIL \*\*\*

COST IN U.S. DOLLARS

FULL ESTIMATED COST

=> s l1 and phenanthridium L2 13 L1 AND PHENANTHRIDIUM

=> dup rem 13
PROCESSING COMPLETED FOR L3

17 DUP REM L3 (0 DUPLICATES REMOVED)

=> s l4 and link?

L5 17 L4 AND LINK?

=> d 15 bib abs 1-17

L5 ANSWER 1 OF 17 USPATFULL on STN

AN 2005:159178 USPATFULL

TI Real-time nucleic acid detection processes and compositions

Rabbani, Elazar, New York, NY, UNITED STATES

Stavrianopoulos, Jannis G., Baysnore, NY, UNITED STATES

Donegan, James J., Long Beach, NY, UNITED STATES Coleman, Jack, East Northport, NY, UNITED STATES

Liu, Dakai, Islip, NY, UNITED STATES

PI US 2005137388 A1 20050623

AI US 2002-96076 A1 20020312 (10)

DT Utility

IN

FS APPLICATION

LREP ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,

CLMN Number of Claims: 542 ECL Exemplary Claim: 1 DRWN 15 Drawing Page(s)

DRWN 15 Drawing Page(s) LN.CNT 6158

AB This invention provides for compositions for use in real time nucleic acid detection processes. Such real time nucleic acid detection processes are carried out with energy transfer elements attached to nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid binding agents. Real time nucleic acid detection allows for the

qualitative or quantitative detection or determination of single-stranded or double-stranded nucleic acids of interest in a sample. Other processes are provided by this invention including processes for removing a portion of a homopolymeric sequence, e.g., poly A sequence or tail, from an analyte or library of analytes. Compositions useful in carrying out such removal processes are also described and provided.

```
ANSWER 2 OF 17 USPATFULL on STN
L5
ΑN
       2005:49898 USPATFULL
       Detection of protein conformations in single cells
TI
       Darzynkiewicz, Zbigniew, Chappague, NY, UNITED STATES
IN
       Traganos, Frank, New York, NY, UNITED STATES
       Juan, Gloria, Sleepy Hollow, NY, UNITED STATES
       Gruenwald, Stefan, Encinitas, CA, UNITED STATES
       US 2005042694
                          Α1
                               20050224
PI
ΑI
       US 2004-954097
                          A1
                               20040929 (10)
RLI
       Continuation of Ser. No. US 1999-256817, filed on 24 Feb 1999, GRANTED,
       Pat. No. US 6821740
PRAI
       US 1998-75908P
                          19980225 (60)
DT
       Utility
FS
       APPLICATION
       DAVID W. HIGHET, VP AND CHIEF IP COUNSEL, BECTON, DICKINSON AND COMPANY,
LREP
       1 BECTON DRIVE, MC 110, FRANKLIN LAKES, NJ, 07417-1880
CLMN
       Number of Claims: 37
ECL
       Exemplary Claim: 1
DRWN
       16 Drawing Page(s)
LN.CNT 2371
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Methods, reagents, and kits are provided that permit flow cytometric
       determination of the phosphorylation status of retinoblastoma
       susceptibility gene protein (pRB) in individual cells. Methods are
       described that permit the hypophosphorylated, active, form of pRB to be
       measured either as an absolute quantity or as a proportion of total
       cellular pRB. Further described are methods that permit pRB
       phosphorylation status to be correlated with cell cycle phase and with
       protein components of the cell cycle. Screening of chemical compounds
       for antiproliferative and antineoplastic activity using the flow
       cytometric assays is demonstrated. Reagent kits that facilitate the
       subject methods are also provided.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L5
     ANSWER 3 OF 17 USPATFULL on STN
AN
       2005:5243. USPATFULL
ΤI
       Novel chemiluminescent reagents
IN
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
       Rabbani, Elazar, New York, NY, UNITED STATES
PA
       Enzo Life Sciences, Inc., New York, NY, 10022 (U.S. corporation)
ΡI
       US 2005004350
                         A1
                               20050106
       US 2004-764388
                               20040123 (10)
ΑI
                          A1
       Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING
RLI
DT
       Utility
      APPLICATION
FS
LREP
       Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
       527 Madison Avenue (9th Floor), New York, NY, 10022-4304
CLMN
       Number of Claims: 17
ECL
       Exemplary Claim: CLM-1-286
DRWN
       15 Drawing Page(s)
LN.CNT 3601
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
       take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin
```

dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays.

They are also applicable to real-time detection processes.

```
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 4 OF 17 USPATFULL on STN
1.5
       2004:321700 USPATFULL
AN
TI
       Labeling reagents comprising aphenylic analogs of rhodamine dyes
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
IN
       Rabbani, Elazar, New York, NY, UNITED STATES
PA
       Enzo Life Sciences, Inc., New York, NY (U.S. corporation)
PΙ
       US 2004254355
                        A1
                              20041216
ΑI
       US 2004-763076
                          A1
                               20040122 (10)
       Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING
RLI
DT
       Utility
       APPLICATION
FS
LREP
       Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
       527 Madison Avenue (9th Floor), New York, NY, 10022-4304
CLMN
       Number of Claims: 286
ECL 
       Exemplary Claim: 1
DRWN
       15 Drawing Page(s)
LN.CNT 4545
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
       take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin
       dyes or composite dyes. These labeling reagents are useful for labeling
```

```
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 5 OF 17 USPATFULL on STN
L5 -
AN
       2004:292946 USPATFULL
ΤI
       Heterodimeric dye composition
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
IN
       Rabban, Elazar, New York, NY, UNITED STATES
       Enzo Life Sciences, Inc., New York, NY, UNITED STATES, 10022 (U.S.
PA
       corporation)
PΙ
       US 2004230036
                          A1
                               20041118
AI.
       US 2004-764389
                          A1
                               20040123 (10)
RLI
       Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING
DT
       Utility
FS
       APPLICATION
LREP
       Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
       527 Madison Avenue (9th Floor), New York, NY, 10022-4304
       Number of Claims: 286
CLMN
```

They are also applicable to real-time detection processes.

probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays.

LN.CNT 4541
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Exemplary Claim: 1

15 Drawing Page(s)

ECL DRWN

L5

This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 6 OF 17 USPATFULL on STN

```
AN 2004:292164 USPATFULL
TI Novel dye labeling composition
IN Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
Rabbani, Elazar, New York, NY, UNITED STATES
PA Enzo Life Sciences, Inc., New York, NY, 10022 (U.S. corporation)
PI US 2004229248 A1 20041118
```

AΙ US 2004-764393 A1 20040123 (10) Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING RLI DT Utility FS APPLICATION Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., LREP 527 Madison Avenue, 9th Floor, New York, NY, 10022-4304 CLMN Number of Claims: 4 ECL Exemplary Claim: CLM-1-286 DRWN 15 Drawing Page(s) LN.CNT 3537 CAS INDEXING IS AVAILABLE FOR THIS PATENT. This invention provides for labeling reagents, labeled targets and AB processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes. CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 7 OF 17 USPATFULL on STN L5 2004:260541 USPATFULL ΑN ΤI Process for preparing novel cyanine dye labeling reagents

Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES TN Rabbam, Elazar, New York, NY, UNITED STATES Enzo Life Sciences, Inc., New York, NY, 10022 (U.S. corporation) PΑ PΤ US 2004203038 A1 20041014 ΑI US 2004-761906 A1 20040121 (10) Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING RLITC Utility FS **APPLICATION** LREP Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,

527 Madison Avenue (9th Floor), New York, NY, 10022-4304

CLMN Number of Claims: 15

ECL Exemplary Claim: CLM-1-286

DRWN 15 Drawing Page(s)

LN.CNT 3584

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

```
L5
     ANSWER 8 OF 17 USPATFULL on STN
ΑN
       2004:248291 USPATFULL
       Process for detecting the presence or quantity of enzymatic activity in
TT
       a sample
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
IN
       Rabbani, Elazar, New York, NY, UNITED STATES
       Enzo Life Sciences, Inc., New York, NY, UNITED STATES, 10022 (U.S.
PA
       corporation)
PΤ
       US 2004192893
                                20040930
                          Α1
ΑI
       US 2004-764417
                          A1
                               20040123 (10)
```

RLI Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING

DT Utility FS APPLICATION

LREP Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., 527 Madison Avenue (9th Floor), New York, NY, 10022-4304

CLMN Number of Claims: 36

ECL Exemplary Claim: CLM-1-286

DRWN 15 Drawing Page(s)

LN.CNT 3665

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 9 OF 17 USPATFULL on STN

AN 2004:228200 USPATFULL

TI Process for detecting the presence or quantity of enzymatic activity in a sample

IN Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES Rabbani, Elazar, New York, NY, UNITED STATES

PA Enzo Life Sciences, Inc., New York, NY, UNITED STATES (U.S. corporation)

US 2004176586 A1 20040909

AI US 2004-764418 A1 20040123 (10)

RLI Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING

DT Utility

PΙ

FS APPLICATION

LREP Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., 527 Madison Avenue (9th Floor), New York, NY, 10022-4304

CLMN Number of Claims: 286 ECL Exemplary Claim: 1 DRWN 15 Drawing Page(s)

LN.CNT 4543

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 10 OF 17 USPATFULL on STN

AN 2004:44526 USPATFULL

TI Characterization of single stranded nucleic acids by melting analysis of secondary structure using double strand-specific nucleic acid dye

IN Wittwer, Carl T., Salt Lake City, UT, UNITED STATES

Dummer, C. Wade, Layton, UT, UNITED STATES

PI US 2004033518 A1 20040219

AI US 2003-423621 A1 20030425 (10)

PRAI US 2002-375640P 20020426 (60)

DT Utility

FS APPLICATION

LREP Richard F. Trecartin, DORSEY & WHITNEY LLP, Suite 3400, Four Embarcadero Center, San Francisco, CA, 94111-4187

CLMN Number of Claims: 52 ECL Exemplary Claim: 1 DRWN 13 Drawing Page(s) LN.CNT 2218

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A novel method for characterizing nucleic acids. A nucleic acid is combined with a double stranded nucleic acid-specific dye to form a detectable complex between the dye and one or more double stranded structures within the nucleic acid. The combination is then exposed to varying temperatures and the fluorescence emission of the dye is measured to determine the melting temperature(s) for the double stranded structures. In some embodiments that melting temperature profile is then compared to melting temperature profiles generated for other nucleic acid(s) to discern differences between the compared nucleic acids.

```
L5
     ANSWER 11 OF 17 USPATFULL on STN
AN
       2003:319498 USPATFULL
       Labeling reagents and labeled targets, target labeling processes and
TT
       other processes for using same in nucleic acid determinations and
       analyses
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
IN
       Rabbani, Elazar, New York, NY, UNITED STATES
ΡI
       US 2003225247
                         A1
                               20031204
ΑI
       US 2002-96075
                          A1
                               20020312 (10)
       Utility
דת
       APPLICATION
FS
       ENZO LIFE SCIENCES, INC., c/o ENZO BIOCHEM, INC., 527 Madison Avenue,
LREP
       9th Floor, New York, NY, 10022
CLMN
       Number of Claims: 286
ECL
       Exemplary Claim: 1
       15 Drawing Page(s)
DRWN
LN.CNT 4499
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
       take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin
       dyes or composite dyes. These labeling reagents are useful for labeling
       probes or targets, including nucleic acids and proteins. These reagents
       can be usefully applied to protein and nucleic acid probe based assays.
       They are also applicable to real-time detection processes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L5
     ANSWER 12 OF 17 USPATFULL on STN
AΝ
       2003:159339 USPATFULL
       FLOW CYTOMETRIC METHODS FOR THE CONCURRENT DETECTION OF DISCRETE
ΤI
       FUNCTIONAL CONFORMATIONS OF PRB IN SINGLE CELLS
       DARZYNKIEWICZ, ZBIGNIEW, CHAPPAQUE, NY, UNITED STATES
IN
       TRAGANOS, FRANK, NEW YORK, NY, UNITED STATES
       JUAN, GLORIA, SLEEPY HOLLOW, NY, UNITED STATES
       GRUENWALD, STEFAN, ENCINITAS, CA, UNITED STATES
PΙ
       US 2003108952
                               20030612
                         A1
       US 6821740
                          B2
                               20041123
       US 1999-256817
AΙ
                         A1
                               19990224 (9)
PRAI
       US 1998-75908P
                          19980225 (60)
DT
       Utility
FS :
       APPLICATION
       SCHNECK & SCHNECK, P.O. BOX 2-E, SAN JOSE, CA, 95109-0005
LREP
CLMN
       Number of Claims: 37
ECL
       Exemplary Claim: 1
DRWN
       16 Drawing Page(s)
LN.CNT 2308
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Methods, reagents, and kits are provided that permit flow cytometric
       determination of the phosphorylation status of retinoblastoma
       susceptibility gene protein (pRB) in individual cells. Methods are
       described that permit the hypophosphorylated, active, form of pRB to be
       measured either as an absolute quantity or as a proportion of total
       cellular pRB. Further described are methods that permit pRB
       phosphorylation status to be correlated with cell cycle phase and with
       protein components of the cell cycle. Screening of chemical compounds
       for antiproliferative and antineoplastic activity using the flow
       cytometric assays is demonstrated. Reagent kits that facilitate the
       subject methods are also provided.
```

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L5 ANSWER 13 OF 17 USPATFULL on STN
- AN 2002:194455 USPATFULL
- TI Multichromophore fluorescent probes using DNA intercalation complexes
- IN Glazer, Alexander N., Orinda, CA, United States

Mathies, Richard A., El Cerrito, CA, United States Peck, Konan, Taipei, TAIWAN, PROVINCE OF CHINA The Regents of the University of California, Berkeley, Berkeley, CA, PA United States (U.S. corporation) PΙ US 6428667 B1 20020806 US 2000-686147 20001010 (9) AΙ Division of Ser. No. US 1997-966398, filed on 7 Nov 1997, now patented, RLI Pat. No. US 6280933 Continuation of Ser. No. US 1993-161231, filed on 2 Dec 1993, now patented, Pat. No. US 5763162 Continuation of Ser. No. US 1992-831823, filed on 6 Feb 1992, now abandoned Continuation-in-part of Ser. No. US 1990-493307, filed on 14 Mar 1990, now abandoned Utility DTGRANTED FS EXNAM Primary Examiner: Whisenant, Ethan C.; Assistant Examiner: Lu, Frank Field, Bret E., Bozicevic, Field & Francis LREP CLMN Number of Claims: 10 ECL Exemplary Claim: 1 0 Drawing Figure(s); 0 Drawing Page(s) DRWN LN.CNT 715 CAS INDEXING IS AVAILABLE FOR THIS PATENT. Novel fluorescent labeling techniques and fluorescent labels are AΒ provided, employing high affinity non-covalently binding and intercalating fluorescent dyes and dsDNA. The dyes find application to provide highly sensitive labeling of nucleic acids in electrophoretic gels and as pre-prepared labels for binding to a wide variety of specific binding pair members. The DNA-dye fluorescer complex can be used for labels in diagnostic assays, detection of specific nucleic acid sequences, and the like. CAS INDEXING IS AVAILABLE FOR THIS PATENT.  $L_5$ ANSWER 14 OF 17 USPATFULL on STN AN2002:69763 USPATFULL ΤI Stabilization of highly sensitive nucleic acid stains in aqueous solutions Wu, Minjie, Thomaston, ME, United States IN White, Hugh W., Camden, ME, United States Kusukawa, Noriko, Salt Lake City, UT, United States Stein, Thomas M., Myersville, MD, United States BioWhittaker Molecular Applications, Inc., Rockland, ME, United States PA (U.S. corporation) \* PΙ US 6365341 B120020402 US 2000-535129 20000324 (9) AΙ DT Utility FS GRANTED EXNAM Primary Examiner: Leary, Louise N. Ratner & Prestia LREP CLMN Number of Claims: 21 ECL Exemplary Claim: 1 DRWN 0 Drawing Figure(s); 0 Drawing Page(s) LN.CNT 358 CAS INDEXING IS AVAILABLE FOR THIS PATENT. AΒ The present invention discloses the use of quaternary compounds as stabilizing agents for highly-sensitive fluorescent nucleic acid stains in aqueous solvents, their use in gels to give increased usable shelf life, and in compositions of solvents, providing ready-to-use stain solutions. CAS INDEXING IS AVAILABLE FOR THIS PATENT. L5 ANSWER 15 OF 17 USPATFULL on STN AN 2001:215176 USPATFULL ΤI Quenching oligonucleotides

Singer, Victoria L., Eugene, OR, United States Haugland, Richard P., Eugene, OR, United States

20011127

20000512 (9)

B1

Molecular Probes, Inc., Eugene, OR, United States (U.S. corporation)

IN

PA

PΙ

ΑI

US 6323337

US 2000-570343

```
PRAI
       US 1999-131782P
                         19990430 (60)
       US 1999-131782P
                          19990403 (60)
       Utility
DT
       GRANTED
FS
EXNAM Primary Examiner: Houtteman, Scott W.
       Helfenstein, Allegra J., Skaugset, Anton E.
LREP
       Number of Claims: 64
CLMN
ECL
       Exemplary Claim: 1
DRWN
       5 Drawing Figure(s); 4 Drawing Page(s)
LN.CNT 1911
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The invention relates to oligonucleotides labeled with an energy
       transfer acceptor useful in conjunction with fluorescent nucleic acid
       stains. The resulting oligonucleotides are useful for decreasing
       background fluorescence during amplification assays and in ligation
       assays, and for detecting hybridization.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 16 OF 17 USPATFULL on STN
1.5
AN
       2001:142076 USPATFULL
TI
       Multichromophore fluorescent probes using DNA intercalation complexes
       Glazer, Alexander N., Orinda, CA, United States
TN
       Mathies, Richard A., El Cerrito, CA, United States
Peck, Konan, Taipei, Taiwan, Province of China
PA
       The Regents of the University of California, Berkeley, CA, United States
       (U.S. corporation)
PΙ
       US 6280933
                          B1
                               20010828
                               19971107 (8)
ΑI
       US 1997-966398
RLI
       Continuation of Ser. No. US 1993-161231, filed on 2 Dec 1993
       Continuation of Ser. No. US 1992-831823, filed on 6 Dec 1992, now
       abandoned Continuation of Ser. No. US 1990-493307, filed on 14 Mar 1990,
       now abandoned
DT
       Utility
       GRANTED
FS
EXNAM Primary Examiner: Campbell, Eggerton A.
       Field, Bret E.Bozicevic, Field & Francis
LREP
       Number of Claims: 19
CLMN
ECL
       Exemplary Claim: 1
DRWN
       No Drawings
LN.CNT 749
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AΒ
       Novel fluorescent labeling techniques and fluorescent labels are
       provided, employing high affinity non-covalently binding and
       intercalating fluorescent dyes and dsDNA. The dyes find application to
       provide highly sensitive labeling of nucleic acids in electrophoretic
       gels and as pre-prepared labels for binding to a wide variety of
       specific binding pair members. The DNA-dye fluorescer complex can be
       used for labels in diagnostic assays, detection of specific nucleic acid
       sequences, and the like.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L5
     ANSWER 17 OF 17 USPATFULL on STN
AN
       1998:64954 USPATFULL
ΤI
       Multichromophore fluorescent DNA intercalation complexes
ΤN
       Glazer, Alexander N., Orinda, CA, United States
       Mathies, Richard A., El Cerrito, CA, United States
       Peck, Konan, Taipei, Taiwan, Province of China
PA
       The Regents of University of California, Berkeley, CA, United States
       (U.S. corporation)
ΡI
       US 5763162
                               19980609
ΑI
       US 1993-161231
                               19931202 (8)
RLI
       Continuation of Ser. No. US 1992-831823, filed on 6 Feb 1992, now
       abandoned which is a continuation-in-part of Ser. No. US 1990-493307,
       filed on 14 Mar 1990, now abandoned
DT
       Utility
FS
       Granted
```

EXNAM Primary Examiner: Campbell, Eggerton A.

LREP Field, BretBozicevic & Reed LLP

CLMN. Number of Claims: 2 ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 672

=>

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Novel fluorescent labeling techniques and fluorescent labels are provided, employing high affinity non-covalently binding and intercalating fluorescent dyes and dsDNA. The dyes find application to provide highly sensitive labeling of nucleic acids in electrophoretic gels and as pre-prepared labels for binding to a wide variety of specific binding pair members. The DNA-dye fluorescer complex can be used for labels in diagnostic assays, detection of specific nucleic acid sequences, and the like.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

```
=> s heterodimer? (4a) dye?
           103 HETERODIMER? (4A) DYE?
=> s 16 not 15
            91 L6 NOT L5
=> s 17 and phenanthridinium
             7 L7 AND PHENANTHRIDINIUM
L8
=> dup rem 18
PROCESSING COMPLETED FOR L8
              5 DUP REM L8 (2 DUPLICATES REMOVED)
=> d 19 bib abs 1-5
     ANSWER 1 OF 5 USPATFULL on STN
L9
       2004:327308 USPATFULL
ΑN
       Methods and compositions for detecting the presence of target nucleic
TI
       acids in a sample
       Kawasaki, Glenn, Seattle, WA, UNITED STATES
IN
       Travis, Bruce M., Seattle, WA, UNITED STATES
                               20041223
PΤ
       US 2004259128
                          Α1
                        A1
ΑI
       US 2004-799925
                               20040311 (10)
                           20031224 (60)
PRAI
       US 2003-532699P
       US 2003-457527P
                           20030324 (60)
DT
       Utility
FS
       APPLICATION
LREP
       BOZICEVIC, FIELD & FRANCIS LLP, 1900 UNIVERSITY AVE, SUITE 200, EAST
       PALO ALTO, CA, 94303
CLMN
       Number of Claims: 31
ECL
       Exemplary Claim: 1
DRWN
       3 Drawing Page(s)
LN.CNT 1585
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Methods and compositions for detecting the presence, e.g.,
       quantitatively, of a target nucleic acid, such as an siRNA, in a sample
       are provided. In the subject methods, a sample is contacted with at
       least two different ligation domains, which may be present on separate
       nucleic acids (e.g., oligonucleotides) or on the same complex, e.g.,
       Combined Oligo, to produce a reaction mixture, where each of the
       different ligation domains includes a domain complementary to a
       different region of the target nucleic acid. The ligation domains of any
       resultant ligation domain/target nucleic acid complexes are then ligated
       to produce a pseudotarget nucleic acid. The presence of any resultant
       pseudotarget nucleic acids in the reaction mixture is then determined in
       order to detect the target nucleic acid in the sample. Also provided are
       systems and kits that find use in practicing the subject methods. The
       subject invention finds use in a variety of applications, including
       therapeutic applications.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 2 OF 5 USPATFULL on STN
L9
       2004:227318 USPATFULL
AN
TI
       Compositions and methods for polynucleotide sequence detection
IN
       Sorge, Joseph A., Wilson, WY, UNITED STATES
       Firmin, Andrew, Jackson, WY, UNITED STATES
PΑ
       Stratagene (U.S. corporation)
       US 2004175704
PΙ
                          A1
                               20040909
ΑI
       US 2003-436231
                         A1
                               20030512 (10)
PRAI
       US 2003-452481P
                          20030306 (60)
DT
       Utility
FS
       APPLICATION
LREP
       PALMER & DODGE, LLP, KATHLEEN M. WILLIAMS / STR, 111 HUNTINGTON AVENUE,
       BOSTON, MA, 02199
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Number of Claims: 61

CLMN

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LN.CNT 2931
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The present invention provides compositions, kits, and methods for
       detecting polynucleotide sequence differences. The method involves
       amplifying a polynucleotide in the presence of a labeled nucleotide
       whose incorporation into the amplified product can indicate the presence
       of a sequence difference within the polynucleotide template. The
       invention is particularly useful for differentiating two or more closely
     related polynucleotide sequences, for example, in determining which
       allele or alleles of a multiallelic organism are present in a target
       polynucleotide.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 3 OF 5 WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN
L9
     2004-055097 [06]
                        WPIDS
ΑN
DNN N2004-044609
                        DNC C2004-022436
     Labeling reagent useful for e.g. determining the amount of nucleic acid in
TΙ
     a sample comprises a marker moiety and a reactive group covalently linked
     together.
DC
     B04 D16 E24 S03
IN
     RABBANI, E; STAVRIANOPOULOS, J G; RABBAM, E; RABBAN, E
PΑ
     (ENZO-N) ENZO LIFE SCI INC; (RABB-I) RABBANI E; (STAV-I) STAVRIANOPOULOS J
     G
CYC
     34
     EP 1348713
PΙ
                     A2 20031001 (200406)* EN 102
         R: AL AT BE BG CH CY CZ DE DK EE ES FI FR GB GR HU IE IT LI LT LU LV
            MC MK NL PT RO SE SI SK TR
     CA 2421552
                    A1 20030912 (200406)
                                           EN
     JP 2004004048
                   A 20040108 (200406)
                                               245
     US 2003225247 A1 20031204 (200406)
     US 2004176586 A1 20040909 (200459)
     US 2004192893 A1 20040930 (200465)
     US 2004203038 A1 20041014 (200468)
     US 2004229248 A1 20041118 (200477)
     US 2004230036
                   A1 20041118 (200477)
     US 2004254355
                    A1 20041216 (200482)
                    A1 20050106 (200504)
     US 2005004350
ADT
     EP 1348713 A2 EP 2003-4894 20030306; CA 2421552 A1 CA 2003-2421552
     20030311; JP 2004004048 A JP 2003-114988 20030311; US 2003225247 A1 US
     2002-96075 20020312; US 2004176586 A1 Div ex US 2002-96075 20020312, US
     2004-764418 20040123; US 2004192893 Al Div ex US 2002-96075 20020312, US
     2004-764417 20040123; US 2004203038 A1 Div ex US 2002-96075 20020312, US
     2004-761906 20040121; US 2004229248 A1 Div ex US 2002-96075 20020312, US
     2004-764393 20040123; US 2004230036 Al Div ex US 2002-96075 20020312, US
     2004-764389 20040123; US 2004254355 Al Div ex US 2002-96075 20020312, US
     2004-763076 20040122; US 2005004350 Al Div ex US 2002-96075 20020312, US
     2004-764388 20040123
PRAI US 2002-96075
                         20020312; US 2004-764418
                                                         20040123;
     US 2004-764417
                          20040123; US 2004-761906
                                                         20040121;
     US 2004-764393
                          20040123; US 2004-764389
                                                         20040123;
     US 2004-763076
                          20040122; US 2004-764388
                                                         20040123
                        WPIDS
AN
     2004-055097 [06]
AB
          1348713 A UPAB: 20040123
     NOVELTY - A labeling reagent (XII) comprises a marker moiety and a
     reactive group covalently linked together.
          DETAILED DESCRIPTION - A labeling reagent of formula (MR) (XII)
     comprises a marker moiety and a reactive group covalently linked together.
          M = marker moiety comprising ligand and/or dye; and
          R = reactive group capable of forming a carbon-carbon linkage with
     the target.
          INDEPENDENT CLAIMS are included for the following:
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(a) a labeled target, labeled by reacting target with (XII) to form a

(b) preparation of cyanine dye labeling reagent of formula (I) involving forming a mixture comprising intermediate compounds of formulae

carbon-carbon linkage between the target and (XII);

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Exemplary Claim: 1

20 Drawing Page(s)

- (Ia) and (Ib), and linking reagents to link (Ia) and (Ib);
- (c) a labeled nucleotide comprising an aphenylic analog of a rhodamine dye, which is attached directly to the nucleotide or indirectly through a linker;
- (d) a heterodimeric dye composition (C1) comprising a dye (a) containing a phenanthridinium moiety and another dye (b) different from (a) and attached through the phenyl ring of the phenanthridinium moiety;
  - (e) determining the amount of nucleic acid in a sample involving:
- (1) forming a mixture of the sample (a dye comprising two phenanthridinium moieties linked through a phenyl group in each of the two moieties, or a dye of formula (IV), or (C1) and reagents for carrying out dye binding, hybridization and/or strand extension) to produce a complex comprising the dye and any nucleic acid present in the sample;
- (2) illuminating the mixture formed at wavelength below 400 nanometer (nm); and
- (3) measuring fluorescent emission from the illuminated mixture, the emission being proportional to the quantity of the nucleic acid present in the sample;
  - (f) a composition comprising at least one of (IV);
  - (g) a chemiluminescent reagent of formula (VIII) or (IX);
- (h) detecting the presence or quantity of enzymatic activity in a sample involving:
- (1) either forming a mixture of the sample, (VIII) or (IX) and reagents and buffers for carrying out chemiluminescent reactions or contacting (VIII) or (IX) and the reagents and buffers with the sample;
- (2) enzymatically converting (VIII) or (IX) into an unstable light-emitting dioxetane form; and
- (3) measuring the quantity of light generated by the enzymatic conversion; and
- (i) a dye composition comprising a compound of formula Rc-Fluorescent  $\ensuremath{\mathsf{Dye}}\,.$
- at least one of R1-R10 = group capable of forming a carbon to carbon bond with a target;

X1, X2 = C, O, N or S;

n = 1-3;

Y = piperidin-1-yl, -NH-(CH2)2-NH-(CH2)2-NH2, N+((CH2)2)-CH2CH2-N+((CH2)2) or N,N-diethyl-N-methylammonium;

Q = (poly)cycloalkyl;

Z = H, aralkyl, alkaryl, (hetero)alkyl, (hetero)aryl, cycloalkyl or cycloheteroalkyl;

R1a and R2a = chemical moieties;

A = cyclic ring;

Ra = chemical linker;

Rb = substrate for non-cleaving enzymatic process;

Rc = unsaturated aliphatic groups, unsaturated heterocyclic groups and/or aromatic groups.

Rla is enzymatically converted into Rlb, which comprises a chemical reactive group Gl. R2a is attached to the cyclic ring through an oxygen atom and comprises a chemical reactive group G2, which reacts with the Gl to convert the dioxetane to an unstable light-emitting dioxetane form. The product of enzymatic process leads to further chemical rearrangement that generate an unstable light emitting dioxetane form. Rc is capable of providing a conjugated system or an electron delocalized system with the fluorescent dye.

USE - For labeling a target; for determining the amount of nucleic acid in a sample; and for detecting the presence or quantity of enzymatic activity in a sample (claimed); and in protein and nucleic acid probe based assays.

Dwg.0/15

TI

PA

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L9 ANSWER 4 OF 5 USPATFULL on STN
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AN 2003:129915 USPATFULL

Method for overcoming bacterial antibiotic resistance

IN Shapiro, Howard M., 283 Highland Ave., West Newton, MA, United States 02465-2513

Shapiro, Howard M., West Newton, MA, United States (U.S. individual)

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PI US 6562785 B1 20030513
AI US 1999-274699 19990323 (9)
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DT . Utility GRANTED

EXNAM Primary Examiner: Low, Christopher S. F.; Assistant Examiner: Mohamed, Abdel A.

LREP Hamilton, Brook, Smith & Reynolds, P.C.

CLMN Number of Claims: 13 ECL Exemplary Claim: 1

DRWN 0 Drawing Figure(s); 0 Drawing Page(s)

LN.CNT 898

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention is drawn to methods of killing bacteria, including antibiotic resistant bacteria, by contacting said bacteria with a membrane permeabilizing compound or combination of compounds and a membrane impermeant toxic agent or combination of agents, resulting in the death of the bacteria without substantial injury to the infected host or patient. The present invention is also drawn to compositions and kits for effecting the method of the present invention. The present invention is further drawn to methods of rendering toxic agents such as toxic organic molecules, membrane impermeant for use in the methods and compositions of the present invention.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L9 ANSWER 5 OF 5 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN DUPLICATE 1
- AN 1994:109442 BIOSIS
- DN PREV199497122442
- TI Heterodimeric DNA-binding dyes designed for energy transfer: Stability and applications of the DNA complexes.
- AU Benson, Scott C.; Mathies, Richard A.; Glazer, Alexander N. [Reprint author]
- CS Dep. Molecular Cell Biology, 229 Stanley Hall, Univ. Calif., Berkeley, CA 94720, USA
- SO Nucleic Acids Research, (1993) Vol. 21, No. 24, pp. 5720-5726. CODEN: NARHAD. ISSN: 0305-1048.
- DT Article
- LA English
- ED Entered STN: 14 Mar 1994
  - Last Updated on STN: 14 Mar 1994
- AB Spectroscopic studies of the complexes of double-stranded (ds) DNA with the polymethylene-amine linked heterodimers thiazole orange-thiazole blue, thiazole orange ethidium, and fluorescein ethidium, in each case show efficient energy transfer from donor to acceptor chromophores (Benson, S.C., Singh, P. and Glazer, A.N. (1993) accompanying manuscript). A quantitative assay of the stability of such complexes during gel electrophoresis is presented. The off-rate of dye from complexes formed at an initial dsDNA bp:dye ratio gtoreq 10:1 follows strict first-order kinetics. The t-0.5 values for the dissociation of a series of related dyes provide a quantitative criterion for the design of DNA-binding fluorophores. Complexes of dsDNA with the monomeric propidium and cyanine dyes, (1-(9-amino-4,7-diazanonyl)-3,8-diamino-6-phenyl-

phenanthridinium bromide trihydrobromide) and (N,N'-tetramethyl-1,3-propanediamino)propyl thiazole orange (4-(3-methyl-2,3-dihydro-(benzo-1,3-thiazole)-2-methylidenyl)-1-(4,4,8-trimethyl-4,8-diazanonyl)-quinolinium diiodide), are much more stable than those with their widely used counterparts, ethidium and thiazole orange. Applications of the new dyes in post-staining of gels and in the multiplex detection of DNA restriction fragments are presented.